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Identification and Validation of Commonly Overexpressed Genes in Solid Tumors by Comparison of Microarray Data

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Abstract

Cancers originating from epithelial cells are the most common malignancles. No common expression profile of solid tumors compared to normal tissues has been described so far. Therefore we were interested if genes differentially expressed in the majority of carcinomas could be identified using bioinformatic methods. Complete data sets were downloaded for carcinomas of the prostate, breast, lung, ovary, colon, pancreas, stomach, bladder, liver, and kidney, and were subjected to an expression analysis using SAM. In each experiment, a gene was scored as differentially expressed if the a value was below 25%. Probe Identifiers were unified by comparing the respective probe sequences to the Unigene build 155 using BlastN. To obtain differentially expressed genes within the set of analyzed carcinomas, the number of experiments in which differential expression was observed was counted. Differential expression was assigned to genes if they were differentially expressed in at least eight experiments of tumors from different origin. The Identified candidate genes ADRM1, EBNA1BP2, FDPS, FOXM1, H2AFX, HDAC3, IRAK1, and YY1 were subjected to further validation. Using this comparative approach, 100 genes were Identified as upregulated and 21 genes as downregulated in the carcinomas. Neoplasia (2004) 6, 744-750

Keywords: Cencer, tumor, microarray, comparative analysis, FOXM1.

Introduction

Solid tumors are the second most common cause of death in the western world. Besides very few successes in rare solid tumors such as testicular cancer, the survival rate of most of these tumors is still low. However, many of these tumors are characterized by inactivating mutations of common tumor-suppressor genes, such as p53, p16, or Rb, and activation of oncogenes such as Her2/heu (1,2.1 Therefore one might guess that not only these genes are differentially expressed within a broad range of tumors, but that there are also other undescribed genes that are differentially expressed in the majority of tumors.

Large-scale gene expression analysis by means of microarrays has yielded large sets of class 2 cancer genes

(3). This enables us to compare the expression profiles of various tumors and to generate sets of common differentially expressed genes. These genes would then represent a pool of interesting candidates used to give new insights into tumor development, and are candidates for new targets for therapy and diagnostics in a variety of cancers. Attempts have been made to compare the gene expression profiles of tumors of a single entity and to assign differentially expressed genes [4–9]. In addition, gene expression differences of tumors from diverse organs have been analyzed with various approaches [10–20].

Gene expression profiles are obtained by different techniques such as DDRT-PCR, SAGE, expressed sequence tag (EST) sequencing, or microarrays [21,22]. Moreover, within the field of microarrays, there are at least three different rival technologies. The most often used techniques, Affymetrix GeneChips and cDNA on glass microarrays, generate different data types and are accompanied by their own challenges within data preprocessing, leading to sets of gene expression data that are not easy to compare [23].

To identify common differentially expressed genes in solid tumors and to overcome these limitations, we have analyzed the differential expression of genes within each experiment by a single method with defined thresholds for 11 different carcinomas. We identified a set of 121 differentially expressed genes and validated eight of these using dot blots containing the transcriptions of different solid cancers.

Materials and Methods

Data Sets

Only data sets containing values for tumors and normal tissues were selected. Data sets for gene expression profiles were obtained by downloading the expression values in tabular form from the Stanford microarray database (http://genome-www.bs.tanford.edu/). GEO (http://www.ncbi.nlm.

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nih.gov/geo/), and the supplementary information of published manuscripts (Table 1). All data sets were used as provided and were normalized if needed by median centering. To unify the used gene identifiers, all probe sequences were compared against the Unigene database version 155 using the BlastN program [24]. Sequences were assigned to a Unigene cluster if their homology was below an e value of e-100, or if their score was higher than 1.5 per base analyzed. Data from the following tumors have been acquired; prostate, breast, lung, ovary, colon, pancreas, stomach, bladder, liver, and kidney.

Identification of Differentially Expressed Genes

All experiments were analyzed using the SAM package (http://www-stat.stanford.edu/~tibs/SAM/index.html) with a cutoff of 25% as a value. To the probes identified by this approach, we assigned the value of 1 if they were overexpressed in the tumor, or -1 if they were underexpressed. If a probe was not differentially expressed, we assigned 0; if it was not analyzed in a given experiment, we assigned a blank field. For all probes, we counted the number of differential expression. To identify genes differentially expressed in the majority of carcinomas, we called only those genes as differentially expressed, which displayed a differential expression within eight of the analyzed 11 cancer entities (73%). For the cancer entities of the kidney, lung, and pröstate, we obtained more than one data set, For these, differentially expressed genes were identified by each experiment as mentioned above. Subsequently, within a type of cancer (e.g., prostate cancer), the differentially expressed genes were compared and scored as differential only if they were identified within at least 50% of the experiments. Thus, for prostate cancer, genes were counted as differentially expressed If they had been identified within at least two experiments—for kidney and lung cancer, if they had been identified within at least one experiment. Such genes were then counted only once for the comparison between the different cancer entities.

Validation of Differential Expression

CDNA clones for ADRM1 (accession no. BM913272). EBNA1BP2 (BU541488), FDPS (BQ877587), FOXM1 (BQ691509), H2AFX (BG757479), HDAC3 (BM468317), IRAK1 (BE250451), and YY1 (BE746736) were obtained from the RZPD (www.rzpd.de) and prepared using the GFX Micro Plasmid Prep kit according to the manufacturer's recommendation (Amersham Pharmacia Biotech, Freiburg, Germany). The identity of the inserts of the obtained clones was confirmed by sequencing of the clones with an ABI 3700 sequencer and BigDye terminators (Applied Biosystems, Weiterstadt, Germany). For the derived sequences, we analyzed their identity against the clones using the program BlastN and the Unigene database. The inserts of the clones were prepared by digesting the DNA with the appropriate restriction enzymes and purification from agarose gels. The fragments were labelled with α-32P-dCTP (~6.000 Ci/mmol) using the Rediprime II kit (Amersham Pharmacia Blotech) and hybridized to a cancer profiling array II (CPA II) using the ExpressHvb hvbridization solution according to the manufacturer's recommendations (Clontech, Heidelberg, Germany).

Autoradiographs were obtained by using a BAS-II scanner (Raytest, Eggenfelden, Germany). For further analysis, intensities were normalized to 8-actin and divided by the normalized intensities of the corresponding normal tissues. These data were subjected to visualization using Treeview (http://rana.lbl.gov/EisenSoftware.htm).

Table 1. Used Gene Expression Profiling Data Sets.

Reference	Tissue	Type of tumor	Type of microarray	Number of tumor	Number of normal	Number of probes/probesets	Source of data
Dyrskjot et al. [30]	Bladder	Carcinoma	Affymetrix	10	4	8,793	GEO
Soflie et al. [19]	Breast	Carcinoma	cDNA	68	4	8,102	SMD
Notterman et al. [18]	Colon	Adenocarcinoma	Affymetrix	18	18	7,129	Supplement
Boer et al. [11]	Kldnev	Renal cell cardnoma	cDNA	32	32	31,500	GEO
Higgins et al. [15]	Kidney	Renal cell carcinoma	cDNA	24	3	22,648	SMD
Chen et al. [12]	Liver	Hepatocellular carcinoma	cDNA	82	75	23,075	SMD
Garber et al. [14]	Lung	Adenocarcinoma and squamous cell carcinoma	cDNA	36 (A) and 13 (S)	5	8,102	SMD
Bhattacharjee et al. [10]	Lung	Adenocarcinoma and squamous cell carcinoma	Affymetrix	127 (A) and 21 (S)	17	12,625	Supplement
Welsh et al. [20]	Ovary	Adenocarcinoma	Affymetrix	27	5	8,793	Supplement
lacobuzio-Donahue et al. [16]	Pancreas	Adenocardnoma	cDNA	11	5	44,500	SMD
Dhanasekaran et al. [13]	Prostate	Adenocarcinoma	cDNA	10	9	9,948	Supplement
Luo et al. [31]	Prostate	Adenocarcinoma	cDNA	16	9	6,500	Supplement
Singh et al. [32]	Prostate	Adenocarcinoma	Affymetrix	52	50	12,625	Supplement
Welsh et al. [33]	Prostate	Adenocarcinoma	Affymetrix	25	9	8,793	Supplement
Leung et al. [17]	Stomach	Adenocardnoma	cDNA	89	23	44,500	SMD

Results

Overlap of Data Sets

A prerequisite for a comparative analysis of gene expression differences is that a high number of genes are analyzed in more than one experiment. This is especially true for comparisons using microarray data sets with incomplete genome representation. However, within the data sets used, the majority of genes is analyzed by at least six experiments, and more than 4000 genes are interrogated by at least eight experiments (Figure 1A).

Identification of Differential Expression

Assignment of significance values is a common procedure in the analysis of gene expression. There are several algorithms known, but the obtained values are rarely corrected for multiple testing. To control the error associated with multiple testing, we used the software SAM (http://wwwstat.stanford.edu/~tibs/SAM/index.html). We tested several cutoffs for the g value generated by SAM in a comparison of four different prostate experiments. From these results, we concluded that a cutoff of 25% would be sensitive enough to generate a list of common differentially expressed genes because we found an overlap of 50% with the data set generated by Rhodes et al. [25] (data not shown). We also tried to use only those genes with a cutoff below 5%, but falled to identify differentially expressed genes in several studies, probably due to a high data variance, which could in part be attributed to a low sample number of normal tissues in different studies (data not shown). When we analyzed the frequency of classification of a gene as differentially expressed, we observed that most of the genes were never classified as differentially expressed and only a minor number were classified more than five times (Figure 1, B and C). Because we were interested to further analyze genes differentially expressed in the majority of Investigated tumors, we chose to examine genes differentially expressed in at least eight tumors. Using this threshold, we identified 100 genes as commonly upregulated and 21 as commonly downregulated in 11 different cancer types from 10 different tissues, as listed in Figure 2A. Within this set, we identified genes such as PCNA and OSF-2, which are well known to be overexpressed within human carcinomas. We then classified the differentially expressed genes according to their purposed function (waw.geneontology.org: Figure 3). However, for most of the identified genes, a function has not assigned yot. Of these genes that were characterized by a category, most of them were grouped into the category of metabosism and cell growth/maintenance. Differences in the distribution of the assigned categories between normal and tumor genes could not be observed. Annotation of the genes revealed that many of the genes are either transcriptional regulators, or take part in the degradation of proteins (Figure 2A).

Validation of Differential Expression

We selected eight upregulated genes based on their degree of differential expression and proposed function (ADRM1, EBNA1BP2, FDPS, FOXM1, H2AFX, HDAC3, IRAK1, and YY1) and investigated their differential expression by hybridization of a gene-specific probe to a CPA II. The CPA II filter contains the transcriptome from 19 different tumor entities. Each tumor is represented by more than one sample, and a corresponding sample of normal tissue from the same donor is provided. For each of the spots on the array, we also obtained the expression values for β-actin as a "housekeeping" control. To evaluate the differential expression of the genes, we divided each value by the corresponding spot value of 6-actin. Furthermore, we divided the normalized values of the tumor by the corresponding normal values (Figure 2B). As expected, most of these genes were overexpressed in a number of samples of different tumor entities displaying the validity of our approach, though we could not observe any upregulation of the genes within the kidney tissues analyzed. Also, only sporadic upregulation was seen in the mammary tumor samples. This might argue against our approach although, except for FOXM1, none of the genes was identified as overexpressed in kidney cancer by our comparative analysis and only five (ADRM1, EBNA1BP2, FDPS, FOXM1, and H2AFX) were identified in

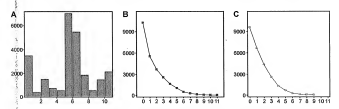


Figure 1. (A) Histogram of the number of genes analyzed by the different experiments. (B and C) Histogram of the number of genes found to be differentially expressed in the different experiments. (B) overexpressed; (C) underexpressed. Harostingly, we did not ligit a gene that is underexpressed in now than nine experiments, including a more interropeneous gene expression in normal fiscure of different ordigis. X-axis: number of experiments; Y-axis: number of genes.

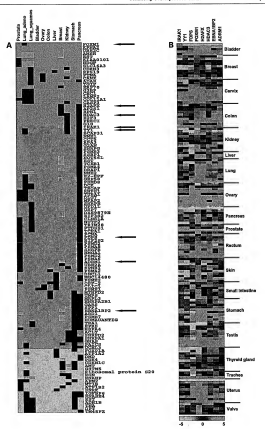


Figure 2. (A) A total of 121 common differentially expressed genes. Red: Overexpressed in tumors; green: underexpressed in tumors; black: not differentially regues z. (27 total of the "Collection Understand y depleased general references and individual section of the community of t

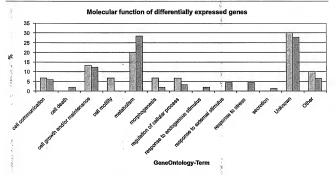


Figure 3. Grouping of identified genes into the molecular function categories of gene ontology using Fatigo (http://fatigo.bioinfo.cnio.es/). Red: Genes overexpressed in tumors; green genes underexpressed in tumors.

breast cancer. Also, we were not able to analyze the quality of the samples in detail. We observed also the heterogeneity of gene expression in the different tumor entities. In our comparative analysis, FOXM1 was overexpressed in 10 of 11 different tumors. This is reflected by the high overexpression of FOXM1 in nearly every tumor analyzed. However, overexpression of FOXM1 could not be observed for some samples of different origins.

Discussion

In an attempt to identify genes commonly overexpressed within solid tumors, we have compared gene expression profiling experiments of 11 different tumors from 10 different organs. We identified the differentially expressed genes using a comparative analysis. The major challenge herein was to unify the different identifiers of the experiments. The data sets used were derived from different microarrays platforms with different structures of identifiers. Whereas Affymetrix GeneChip provides a probeset identifier with a target sequence, the obtained data sets from cDNA microarrays provide only an accession number of the used clones. This leads to the need for a large computational effort. Unfortunately, most of the gene expression data have not been published as raw data so far. Therefore, published articles are the prevalent repository for differentially expressed genes. Comparison of these data has its own merit [26], but only analysis of the raw data circumvents the inherent problems of microarrays within the field of data normalization and expression level assignment. Moreover, comparison of published data is hampered by different modes of gene expression analysis, possibly leading to a subjective choice of genes.

Within this set of analyzed genes, most of them have never been assigned as differentially expressed. Also, only a few genes are identified as differentially expressed in more than five experiments (Figure 1, B and C). From those data, we might conclude that a comparison of microarray data leads not to an accumulation of false positives but to subsets of genes that might be worthwhile in further analyses.

We were interested in those whose expression is deregulated in a majority of tumors. Therefore, we decided to concentrate on genes that are differentially expressed in 8 of 11 analyzed tumors. We identified 100 genes as commonly overexpressed and 21 as commonly downrequlated within carcinomas investigated. Interestingly, we were able to identify more genes as commonly overexpressed than genes commonly underexpressed in tumors. This might be due to a higher variance of the observed gene expression in normal tissues. This variance might result from an inadequate low number of these tissues in individual comparisons (i.e., there were only 4 normal breast tissues and 68 tumor tissues analyzed by Sorlie et al. [19]) (Table 1). As normal tissue has to abide by the precise function of an organ, which is different between organs, a tumor has, first of all, the need to grow. To accomplish this task, a tumor has to circumvent several safeguard functions of cell maintenance and growth limitations [1]. To bypass these, only a few genes have to be modified, which might lead to a common tumor phenotype. Therefore, gene expression of tumors of different organs might probably be more homogenous than the gene expression of normal tissues from the same organs.

Within the set of overexpressed genes, we identified genes from different functional categories. Interestingly,

genes from the protein degradation pathway by the ubiquitin ligation/proteasome pathway as well as transcription factors were often found. Those genes have already been described as overexpressed within tumors [1,27], therefore demonstrating the feasibility of our approach. In addition, these genes represent key members for new targets for cancer therapy. Validation of differential expression is a key for successful microarray experiments. This should also refer to comparative or meta-analysis. However, tissue samples of different tumor entities with an adequate number of samples can rarely be obtained. We validated the differential expression of 8 of 100 overexpressed genes and observed an overexpression in most cases. That not all genes were overexpressed in all samples might be attributed to either the heterogeneity of gene expression in tumors or perhaps errors of the sampling of tissues, or that the gene was falsely assigned as differentially expressed by our method.

Within the group of overexpressed genes, we identified FOXM1 and ADAR1. FOXM1 is activated by the hedgehog signailing pathway [28], indicating a common activation of this elemental developmental pathway in cancers. ADAR is a member of a diverse family of proteins involved in the editing of mRNA. Overexpression of this gene might lead to a higher amount of edited RNA and might therefore lead to mutated proteins. A high level of ADAR within cancers therefore might act as a mutator mutation. That enzymes of the RNA editing pathway are capable of acting in this manner has been shown for APOBEC [29].

in conclusion, using a novel approach to compare gene expression data, we identified a set of genes that might be useful in the further analysis of fundamental signal transduction pathways that lead to carcinomas.

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